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Polyorganosiloxane series thermoplastic resin and composition thereof.

We A polyorganosiloxane series thermoplastic resin comprising a graft copolymer obtained from a virty moreomer and a modified polyorganosiloxane, and a thermoplastic resin composition containing the same. The presin and the composition thereof have excellent sliddbilly, abrasion resistance, weather resistance, cold resistance and impact resistance, and can be applied to new fields such as sliding parts, parts for cold district, outdoor parts, alc.

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## POLYORGANOSII OXANE SERIES THERMOPLASTIC RESIN AND COMPOSITION THEREOF

#### FIELD OF THE INVENTION

This invention relates to a polyorganositoxane series thermoplastic resin containing efficiently graft polymerized viry! monomer(s) and having remarkably excellent slidability, abrasion resistance, weather esistance, ood resistance, and impact resistance.

#### BACKGROUND OF THE INVENTION

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Hitherto, for improving the impact strength of a thermoplastic resin, a technique of modifying the resin by rubber has been established.

For example, there are an ABS resin obtained by reinforcing a styrene-acrylonitrile resin (AS resin) with a butadiene rubber and an AAS resin obtained by reinforcing as AS resin with an acrylic rubber.

As such a base rubber component, a silicone rubber is considered but when polyorganosiloxane is simply compounded with a thermoplastic resin. the compounded product is insufficient in impact resistance since these components are interior in compatibility with each other. Thus, a technique of grafting a vinyl monomer on a rubber as in an ABS resin is required, but a polyorganosiloxane is generally poor in a reactivity with a vinyl monomer and hence it is not easy to form a graft polymer of a polyorganosiloxane. Thus, for forming the craft conditivers of this kind, several methods are proposed.

For example, it is proposed in JP-A-50-109282 (the term "JP-A" as used herein means an "unexamıned published Japanese patent application") that a graft copolymer having an improved impact strength is obtained by polymerizing a vinyl monomer in the presence of a polyorganosiloxane having a vinyl group or an allyl droup.

A method of improving the impact strength of a vinyl polymer by using a polyorganosiloxane having a mercapto group in place of the polyorganosiloxane having a vinyl group or an allyl group is also proposed in JP-A-52-130885.

Furthermore, it is proposed in JP-A-60-252613, JP-A-61-106614, and JP-A-61-136510 that a graft copolymer having an excellent impact strength is obtained with a high graft efficiency by polymerizing a winyl monomer in an emulsion of a polyorganosiloxane having an acryl group or a methacryl group.

However, when the polyorganosiloxane having a vinyl group or an allyl group or the polyorganosiloxane having a mercaplo group is used, the polyorganosiloxane is poor in a reactivity with a vinyl monomer. Accordingly, an apparent graft ratio calculated from an amount of a gel formed, that is, a ratio of the grafted vinyl polymer to the amount of the polyorganosiloxane, is small.

Therefore, an interface adhesive strength between such a polyorganosiloxane and a vinyl polymer is low, thereby causing a emarkable delarnination, whereby there is a problem that a graft copolymer having good appearance and a sufficient impact strength cannot be obtained.

Also, when the polyogenosifloxane having an acryl group or a methacryl group is used, a graft copolymer having an improved impact strength is obtained, but the heat stability thereol is poor and the graft ratio is greatly lowered by the high-temperature molding and the relenting during molding. Consequently, the impact strength, appearance, and luster of the moldings are greatly reduced.

## SUMMARY OF THE INVENTION

The present invention has been made under such circumstances of the conventional techniques.

Accordingly, one object of this invention is to provide a polyorganosiloxane series thermoplastic resin having excellent weather resistance, cold resistance, slidability, and abrasion resistance with an excellent graft reactivity.

Another object of this invention is to provide a thermoplastic resin composition comprising the polyorganosiloxane series thermoplastic resin and other thermoplastic resin(s).

The polyorganositoxane series thermoplastic resin according to this invention (hereinafter sometimes referred to as simply "the thermoplastic resin") comprises a graft copolymer (V) (hereinafter sometimes

referred to as "component (V)") obtained by grafting at least one kind of vinyl monomer (IV) (hereinafter sometimes referred to as "component (IV)") onto a modified polyorganosiloxane (III) (hereinafter sometimes referred to as "component (III)"), the component (III) being obtained by condensing from 90 to 99.8% by weight of organosiloxane (I) thereinafter sometimes referred to as "component (I)") having a structural unit 5 represented by the formula (A)

R',SiO(4-n) 2 (A)

wherein R' represents a substituted or unsubstituted monovalent hydrocarbon group and n represents 0 or an integer of from 1 to 3, and from 10 to 0.2% by weight of a graft crosslinking agent (II) (hereinafter sometimes referred to as "component (II)") having both an unsaturated group represented by the following 10 formula (B)

$$CH_2 = C$$
(B)

wherein R2 represents a hydrogen atom or an alkyl group having from 1 to 6 carbon atoms, and an alkoxysilyl group, the sum of component (I) and component (II) being 100% by weight.

## DETAILED DESCRIPTION OF THE INVENTION

The organosiloxane (I) used in this invention has the structural unit shown by the above-described formula (A) and has a straight chain, branched or cyclic structure, and is preferably the organosiloxane having a cyclic structure.

In formula (A) described above. R' represents a substituted or unsubstituted monovalent hydrocarbon group as described above, and examples of the monovalent hydrocarbon group are a methyl group, an ethyl group, a propyl group, a vinyl group, a phenyl group and those groups each substituted with a halogen atom or a cyano group.

Further, in the formula (A), n is 0 or an integer of from 1 to 3.

Specific examples of the organosiloxane (I) are cyclic organosiloxanes such as hexamethylcyclotrisiloxane, octamethylcyclotetrasiloxane, decamethylcyclopentasiloxane, dodecamethylcyclohexasiloxane. trimethyltriphenylcyclotrisiloxane, etc., and straight chain and branched organosiloxanes.

In addition, the organosiloxane (I) may be a previously condensed polyorganosiloxane having a weight average molecular weight of from about 500 to 10,000 calculated as a polystyrene.

When the organosiloxane (I) is a polyorganosiloxane, the terminal of the molecular chain may be blocked by, e.g., a hydroxy group, an alkoxy group, a trimethylsilyl group, a dimethylvinylsilyl group, a methylphenylvinylsilyl group, a methyldiphenylsilyl group, etc.

The graft crosslinking agent (II) used in this invention is a compound having the unsaturated group shown by the formula (B) described above and an alkoxysilyl group.

In the formula (B),  $\mathbb{R}^2$  is a hydrogen atom or an alkyl group having from 1 to 6 carbon atoms. Preferred of those is a hydrogen atom or an alkyl group having from 1 or 2 carbon atoms, and more preferred is a 45 hydrogen atom or a methyl group.

Specific examples of the graft crosslinking agent are p-vinylphenylmethyldimethoxysilane, 1-(m-vinylphenyl)methyldimethylisopropoxysilane. 2-(p-vinylphenyl)ethylmethyldimethoxysilane. 3-(p-vinylphenoxy)propylmethyldiethoxysilane, 3-(p-vinylbenzoyloxy)propylmethyldimethoxysilane, 1-(o-vinylphenyl)-1,1,2-1-(p-vinylphenyl)-1,1-diphenyl-3-ethyl-3,3-diethoxydisiloxane, trimethyl-2,2-dimethoxydisilane. [3-(p-isopropenylbenzoylamino)propyl]-50 vinylphenyl[3-(triethoxysilyI)propyI]diphenylsilane,

phenyldipropoxysilane and mixtures thereof.

vinylbenzoyloxy)propylmethyldimethoxysilane are preferred, and p-vinylphenylmethyldimethoxysilane is more preferred.

The amount of the graft crosslinking agent, component (II), used is from 0.2 to 10% by weight, and preferably from 0.2 to 5% by weight, based on the sum of component (I) and component (II). If the amount thereof is less than 0.2% by weight of the sum, a high graft ratio is not obtained in the graft polymerization of the modified polyorganosiloxane (III) and the vinyl monomer (IV). As a result, the interface adhesive strength between the modified polyorganosiloxane (IIII) and the vinyl polymer grafted thereto is decreased, thereby causing detarmination, and the graft copolymer (V) having a sufficient impact strength cannot be obtained

On the other hand, if the amount of the graft crosslinking agent (II) is more than 10% by weight of the 5 sum, the graft ratio is increased, but the degree of polymerzation of the grafted vinyl polymer is reduced with the increase of the graft crosslinking agent (II) to reduce the molecular weight of the vinyl polymer, and as a result, a sufficient impact strength cannot be obtained.

On the other hand, the graft copolymer to a silicone polymer tends to detenorate by light oxidation up to a deep portion of the grafted vinyl polymer due to the high light transmittance and high gas permeability (oxyqeno for silicona.

To overcome these problems, a function as a UV absorbing layer may be imparted to the silicone by adding an ultravolet (UV) absorber.

However, if the UV absorber is simply added, the existence of the UV absorber is predominated in the set of the wnyl polymer having a high compatibility with the UV absorber since there is a difference in 5 compatibility of the UV absorber for the silocane and the vinyl polymer, and a sufficient effect as the UV absorbing layer for the silocone cannot be obtained.

To overcome this problem, it is preferred to chemically combine a UV absorber with the siloxane and in this case, it is preferred to introduce a group containing a UV absorbing group into the siloxane as R1 in the formula (A) described above.

The group having the UV absorbing group includes a group represented by the following formula

$$\begin{array}{c|c}
 & x & \overline{z} \\
\hline
 & x & \overline{Q} \\
\hline
 & Q & CH_2 \gamma_n
\end{array}$$

wherein X represents

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Y represents H or OH; Z represents H, OH, or OR<sup>2</sup>; and n represents an integer of from 1 to 5 (R<sup>2</sup> described above is an alkyl group having 1 to 18 carbon atoms); with proviso that when Y is H, at least one Z is OH.

More practically, the group shown by the following formula is preferred.

The content of the group having the UV absorbing group is from 0.02 to 10% by weight, and preferably form 0.1 to 5% by weight, based on the total amount of the organic groups bonded to the silicon atom. If the content is less than 0.02% by weight, the effect in ultraviolet resistance is insufficient, while if the so content is more than 10% by weight, the heat resistance is reduced in the case of using as a composite material by blending the thermole-stic resin with other organic polymer.

Specific examples of the organic stiticon compound (VI) (hereinafter sometimes referred to as "component (VI)") having such a UV absorbing group are

and the following organic silicon compound is preferred

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In the case of forming the polyorganosiloxane as component (III) by polycondensing the organosiloxane (an organosiloxane (i) having the structual unit represented by the following formula (A):

R<sup>1</sup>/<sub>2</sub>Si<sub>4/2</sub> (A).

R<sup>2</sup>/<sub>2</sub>Si<sub>4/2</sub> (A)

Wherein A? Represents a substituted or unsubstituted monovalent hydrocarbon group and n represents O or an integer of from 1 to 3 (hereinafter sometimes referred to as "component (i)"), the organosilicon compound (component (II)) and the UV absorbing group-containing organosilicon component (III) and component (IV) are component (IV) and the UV absorbing group in component (IV) become from 0.02% to 10% of the total amount of the organic groups bonded to the silicon atom of component (III).

The compounding of these components is conducted, for example, such that they are compounded in the amounts of from 80 to 99,6% by weight of the organosiloxane as component (1), from 10 to 0.2% by weight, preferably from 7 to 0.5% by weight, and more preferably from 5 to 1% by weight, preferably from 7 to 0.5% by weight, and more preferably from 7 to 0.5% by weight, and more preferably from 7 to 0.5% by weight, and more preferably from 7 to 0.5% by weight, and more preferably from 5 to 1% by weight, of the UV absorbing group-having organic silicon compound as component (10), the sum of component (1), component (10) and component (10) being 100% by weight.

By polycondensing the above components, the desired polyorganosiloxane can be obtained. In this invention, the modified organosiloxane (III) can be produced by mixing under shearing the organosiloxane (I) and the graft crosslinking agent (II) In the presence of an emulsifying agent such as an adjybranzensufforic acid, etc., using a homomiser, and condensing them. The emulsifying agent functions as an emulsifying agent for the organosiloxane (I) and also as a condensation initiator.

Examples of the emulsifying agent are an alighatic substituted naphtalenesulfonic acid, an alighatic Examples of the emulsifying acid, and alighatic substituted diphenyl ether sulfonic acid, each alphatic group having a carbon lenoth of from 6 to 18 carbon atoms.

Further, nonionic emulsifying agnets such as polyoxyethylene alkyl ester, polyoxyethylene alkylaryl 5 eiher, etc., can be artifact.

The amount of the emulsifying agent used is usually from 0.1 to 5% by weight, and preferably from about 0.3 to 3% by weight, based on the sum of component (I) and component (II).

n addition, water is used in the production of the organosiloxans, and the amount of water is usually from 100 to 500 parts by weight, and preferably from 200 to 400 parts by weight, per 100 parts by weight of the sum of component (j) and component (ii).

The condensation temperature is usually from 5 to 100°C.

In addition, in the case of producing the modified polyorganosiloxane (III), a crossinking agent can be added to the system as the third component for improving the impact resistance of the resis obtained. Examples of the crossinking agent are infunctional crosslinking agents such as methyltrimethoxyslane, ethyltrinethoxyslane, ethyltrinethoxyslane,

In addition, the weight average motecular (calculated as a polystyrene) of the modified polyorganosiloxane (III) thus obtained is usually from 10,000 to 5,000,000, preferably from 30,000 to 1,000,000, and more preferably from 50,000 to 3,000,000.

By graft polymerizing the vinyl monomer (IV) onto the modified polyorganosiloxane (III) thus obtained, the polyorganosiloxane series thermoplastic resin of this invention containing the graft copolymer (V) is obtained.

Examples of the vinyl monomer (IV) which is used to obtain the thermoplastic resin of this invention are aromatic alkenyl compounds such as styrene, e-methylstyrene, sodium styrenesulfonate, etc.; methacrylic add seters such as methyl methacrylate, ethyl methacrylate, etc.) when the methacrylate, butyl methacrylate, allyl methacrylate, hydroxyethyl methacrylate, dimethylaminoethyl methacrylate, etc.; acrylic acid esters such as methyl scrylate, ethyl acrylate, butyl acrylate, etc.; vinyl cyanide compounds such as acrylinitine, methacrylorithie, etc. olefins such as ethylene, propylene, etc.; conjugated diolefins such as butadiene, isoprene, chloroprene, etc.; and vinyl acetate, vinyl chloride, vinylidene chloride, triallyl isocyanurate, acrylic acid, methacrylic acid. N-phenylmaleimide, N-cyclohexylmaleimide, maleic anhydnde, etc. Those may be used alone or as mixture therace!

The graft copolymer (V) containing from 65 to 75% by weight styrene and from 35 to 25% by weight acrylonitrile as the winyl monomers (IV) is preferred for improving the impact resistance of the polyor-ganosiloxane series thermoplastic resist not this invention.

In addition, in the case of graft polymerizing the vinyl monomer (IV) onto the modified polyorganositoxane (III), the proportion of the component (III) is from 5 to 80% by weight, and preferably from 10 to 60%
by weight, and the proportion of the component (IV) is from 95 to 20% by weight, and preferably from 90 to
40% by weight, the sum of the component (III) and the component (IV) being 100% by weight.

If the proportion of the component (III) is less than 5% by weight, a sufficient impact strength cannot be obtained, while if the proportion of the component (III) is more than 80% by weight, the proportion of the winyl polymer to be graft bonded is reduced, thereby a sufficient interlayer adhesive strength cannot be obtained between the modified polyporganosiloxane (III) and the winyl polymer, and as a result, the thermoplastic resin obtained is Interfor in appearance and impact strength.

The graft ratio of the graft copolymer (V) thus obtained is usually about 20% by weight or more, preferably about 80% by weight or more, and more preferably about 100% by weight or more.

When the graft ratio of the graft copolymer (V) is high as described above, the interface adhesive strength between the graft copolymer and the vinyl polymer which was not directly grafted thereto is increased, and the modified polyorganositoxane (III) is uniformly dispersed in the vinyl polymer to provide the thermoplastic resin having good appearance and an excellent impact strength.

The thermoplastic resin of this invention further contains, in addition to the graft copolymer (V) thus obtained, a vinyl polymer which is an ungrafted polymer of the vinyl monomer (IV), and the content of the graft copolymer (V) in the thermoplastic resin is usually 5% by weight or more, and preferably 10% by weight or more.

In the production of the thermoplastic resin of this invention, the vinyl monomer (IV) is graft-polymerized to the modified polygraposiloxane (III) by a conventional radical polymerization to provide a composition containing the graft cooolymer (\cappa)

In this case, according to the kind of a radical polymerization initiator used, it is necessary to neutralize with alkali a latex of the modified polyorganosiloxane which is acidic by the alkylbenzenesulfonic acid as described above. Examples of the alkali for neutralizing the acid latex are sodium hydroxide, polassium hydroxide, sodium carbonate, sodium hydrogencarbonate, Inethanolamine, thethylamine, etc.

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The amount of the radical polymerization initiator used is usually from 0.05 to 5 parts by weight, and preferably from 0.1 to 3 parts by weight, per 100 parts by weight of the vinyt monomer (IV).

s In this case, the radical polymenzation is preferbly carried out by an emulsion polymerization or a solution polymenzation.

In conducting the emulsion polymerization, a conventional emulsifying agent, the above-described radical polymerization initiator, a chain transfer agent, etc., are used.

Examples of the emulsylving agent are amonic emulsifying agents such as sodium dodecyfbenzenesul20 fonate, sodium laurytsulfate, sodium diphenyl ether disolfonate, succinc acid dialkali ester sodium sulfonate, 
etc.: and nonionic emulsifying agents such as polyoxyethylene alkyl ester, polyoxyethylene alkylaryl ether, 
etc. They can be used alone or as a combination thereof.

The amount of the emulsifying agent is usually from about 0.5 to 5% by weight based on the weight of the virul monomer (IV)

Examples of the chain transfer agent are mercaptans such as t-dodecy/mercaptan, octhy/mercaptan, nletradecy/mercaptan, n-nexy/mercaptan, etc., and halogen compounds such as carbon tetrachloride, etby/lene bromide, etc.

The amount of the chain trasfer agent is usually from 0.02 to 1% by weight based on the weight of the vinyl monomer (IV).

The emulsion polymerization is carried out at a polymerization temperature of from 5 °C to 100 °C, and preferably from 50 °C to 90 °C. for a polymenization time of from 0.1 hour to 10 hours using the radical polymenization initiator, the emulsifying agent and the chain transfer agent in the amounts defined above together with water in an amount of usually from 100 to 500 parts by weight to 100 parts by weight of the vinyl monomer (IV) and, if necessary, an electrolyte, a pH controlling agent, etc.

In addition, the emulsion polymerization can be practiced by adding the vinyl monomer (IV) and the radical polymerization initiator to the latex containing the modified polyorganosiloxane (III) obtained by the condensation of the organosiloxane (I) and the graft crosslinking agent (II).

On the other hand, in the case of the solution polymerization, the modified polyorganosiloxane (III) and the vinyl monomer (IV) are dissolved in an organic solvent and the radical polymerization is conducted by adding the

Examples of the organic solvent which is used for the solution polymenization are toluene, n-hexane, cyclohexane, chloroform, tetrahydrofuran, etc.

The solution polymerization is carried out at a polymerization temperature of from 5 °C to 150° C, and preferably from 50° C to 130° C, for a polymerization time of from 1 hour to 10 hours using the radical preferably from 50° C to 130° C, for a polymerization time of from 1 hour to 10 hours using the radical solvent preferably from 50° C to 130° C, for a polymerization initiator and, if necessary, the chain transfer agent, etc., in the amounts as defined above together with, usually, an organic solvent in an amount of from 80 to 500 parts by weight per 100 parts by weight of the vinyl monomer (IV).

In the case of the solution polymerization, the formation of impurities is greatly reduced as compared to the case of the emulsion polymerization.

In the case of producing the polyorganosiloxane series thermoplastic resin of this invention by the emulsion polymerization, the product is usually coagulated by a salting out method and the powdery product thus obtained is purified by washing with water, followed by drying.

In the case of the solution polymerization, unreacted monomers and the solvent are removed from the reaction mixture by a steam distillation and the massive resin obtained is finely ground and dried for purification of the norm-time.

The thermoplastic resin of this invention containing the graft copolymer (IV) obtained by each method described above can be formed into pellets by a kneading machine such as an extruder, etc.

In this case, according to the required performance, other conventional polymer can be properly

blended with the thermoplastic resin obtained in an amount of about 99% by weight or less, and preferably about 90% by weight or less, to pronde a polyorganosiloxane series thermoplastic resin composition (this blend is also referred to as "thermoplastic resin composition").

Examples of the conventional polymer which can be used with the thermoplastic resin of this invention are diene series rubbers such as polybutadiene, a butadiene-styrene copolymer, an acrylonitrile-butadiene copolymer, polyisoprene, natural rubbers, etc.; olefinic rubbers such as acrylic rubber, a ethylene-propylene copolymer, an ethylene-propylene-diene copolymer, chlonnated butyl rubber, chlorinated polyethylene, etc.: aromatic vinyl conjugated diene series block copolymers such as a styrene-butadiene block copolymer. a styrene-butadiene-styrene block copolymer, a styrene-butadiene-styrene radial tereblock copolymer, etc.; 10 hydrogenated products of those block copolymers; polypropylene, polyethylene, polystyrene, a styreneacrylonitnie copolymer, rubber-reinforced polystyrene (HIPS), an acrylonitrile-butadiene-styrene resin (ABS resin), an acrylonitrile-ethylenepropylene-styrene resin (AES resin), a methyl methacrylate-butadiene-styrene resin (MBS resin), an acrylonitrile- butadiene-methyl methacrylate-styrene resin, an acrylonitrile-n-butyl acrylate-styrene resin (AAS resin), polyvinyl chlonde, polycarbonate, polyethylene terephthalate, polybutylene terephthalate, polyacetal, polyamide, an epoxy resin, polyvinylidene fluoride, polysulfone, and ethylene-vinyl acetate copolymer, a PPS resin, polyether ether ketone, a PPO resin, polyacrylate, a styrenemethyl methacrylate copolymer, a styrene-maleic anhydride copolymer, a rubber-modified PPO resin, a styrene-maleimide series copolymer, a rubber-modified styrene-maleimide copolymer, a polyamide series elastomer, and a polyester series elastomer.

The pellets of the thermoplastic resin (composition) are molded by a conventional means such as compression molding, injection molding, etc.

In the thermoplastic resin composition comprising the thermoplastic resin containing the graft copolymer (V) and other polymer as described above, by using the wnyl monomer having a carboxyl group or an epoxy group and other vinyl monomer as the graft monomers to the graft copolymer (V), the thermoplastic resin shows an excellent compatibility with other polymer used.

In particular, when the thermoplastic resin composition is formed using a polymer having a polarity, such as polyamide and polyester, as the polymer, the thermoplastic resin and the polymer are excellent in compatibility with each other and the composition having excellent physical properties can be obtained.

Examples of the carboxyl group-containing unsaturated vinyl monomer are acrylic acid, methacrylic acid, crotonic acid, cinnamic acid, lateconic acid, analeic acid, and acid anhydrides such as maleic anhydride. etc. Of those, acrylic acid and methacrylic acid are preferred.

Examples of the epoxy group-containing vinyl monomer are glycidyl methacrylate. glycidyl acrylate. vinyl glycidyl ether, allyl glycidyl ether, a glycidyl ether of hydroxyalkyl (meth)acrylate, a glycidyl ether of polyalkyleneglycol (meth)acrylate, and glycidyl itaconate.

Examples of the other viny! monomer which is used together with the carboxyl group- or epoxy groupcontaining viny! monomer as the graft monomers to the graft copolymer (V) are aromatic alkeny!
compounds such as styrene, o-methylstyrene, vinybytene, monochlorostyrene, dichlorostyrene, monochromostyrene, dichlorostyrene, p-butylstyrene, ethylstyrene, vinyhaphthalene, o-methylstyrene, p-methylstyrene, dimethylstyrene, sodium styrenesulfonate, etc.: methacrylic acid esters such as methyl
methacrylate, ethyl methacrylate, 2-ethylhexyl methacrylate, butyl methacrylate, lallyl methacrylate, hydroxyethyl methacrylate, ethyl methacrylate, etc.: avrighte, acid esters such as methyl
ethyl acrylate, butyl acrylate, etc.: vinyl cyanide compounds such as acytonitnie, methacrylonitrilic, etc.:
etc.; vinyl acetate, vinyl clohoide, vinyllidene chloride, triallyl isocyanurate, and maleimides such as N
ethylmaleimide, N-methylmaleimide, N-echlorophenylmaleimide, N-cyclohexylmaleimide, etc. They may be used allone or as a mixture thereof.

In the case of graft polymentzing the carboxyl group-containing or epoxy group-containing viryl monomer and other viryl monomer sends polymer, the proportion of the polyorganosiloxane series polymer is from 5 to 90% by weight, preferably from 10 to 70% by weight, and so more preferably from 20 to 65% by weight, and the sum of the proportions of the viryl monomers is from 95 to 10% by weight, and more preferably from 90 to 30% by weight, and more preferably from 80 to 40% by weight.

If the proportion of the polyorganositoxane series polymer is less than 5% by weight, a sufficient impact strength is not obtained, while if the proportion thereof is more than 90% by weight, a preferred powder of the graft copolymer is not obtained and the powder is not dispersed at blending the copolymer.

In addition, the content of the polyorganositoxane in the whole composition is preferably from 5 to 30% by weight, if the content is less than 5% by weight, a sufficient impact strength is not obtained, while if the content is more than 30% by weight, the heat resistance is sometimes reduced.

The content of the carboxyl group-containing vinyl monomer and/or the epoxy group-containing vinyl

monomer in the graft copolymer is preferably from 0.05 to 30% by weight, and more preferably from 0.1 to 20% by weight. If the content is less than 0.05% by weight, the compatibility increasing effect for the graft copolymer with a resin such as polyphenylene sulfide, etc., is decreased, while if the content is more than 30% by weight, the degree of gelation is increased and the extrusion moldability of the thermoplastic resin 5 (composition) obtained becomes sometimes difficult. Other viryl monomer described above is not an essential component in this invention, but from the relationship between the amount of the organosiloxane series copolymer in the graft polymer and the amount of the carboxyl group-containing or epoxy-containing vinyl monomer, it is preferred to use other vinyl monomer in the amount of giving the above-described proportion of the carboxyl group-containing or epoxy group-containing vinyl monomer in the graft polymer.

In particular, the weight ratio of the carboxyl group-containing vinyl monomer/other vinyl monomer is preferably from 0.5.99.5 to 95.5 and more preferably from 1.99 to 90.10.

Furthermore, the content of the carboxyl group-containing vinyl monomer and/or the epoxy groupcontaining vinyl monomer in the whole component is preferably from 0.01 to 5% by weight, more preferably from 0.02 to 4% by weight, and particularly preferably from 0.1 to 3% by weight. If the content thereof is less than 0.01% by weight, the compatibility increasing effect for the graft copolymer with, for example, a polyphenylene sulfide series resin is reduced, while if the content is more than 5% by weight, the degree of gelation is increased and extrusion molding becomes sometimes unapplicable.

Furthermore, the graft ratio of the vinyl monomer to the polygrganosiloxane series copolymer in the graft copolymer is preferably 10% by weight or more. If the graft ratio is less than 10% by weight, the 20 appearance of the molded product is sometimes undesirably reduced.

The proportion of the graft copolymer (A) in the thermoplastic resin composition of this invention is from 5 to 90% by weight, preferably from 10 to 80% by weight, and more preferably from 15 to 70% by weight. If the proportion thereof is less than 5% by weight, the impact resistance of the thermoplastic resin composition obtained is reduced, while if the proportion is more than 90% by weight, the heat resistance and the chemical resistance of the composition are reduced.

When the graft copolymer obtained by graft polymerizing carboxyl group- or epoxy group-containing virill monomer to the polyorganosiloxane series polymer is used, the preferred thermoplastic resin compositions containing the graft copolymer and other polymer are as follows.

A thermoplastic resin composition comprising

(A) from 5 to 90% by weight, and preferably from 10 to 80% by weight, of the graft copolymer obtained by graft cooplymenizing from 95 to 10% by weight of sum of the epoxy group-containing vinyl monomer and the other vinyl monomer to from 5 to 90% by weight of the modified polyorganosiloxane (III)

(8) from 10 to 95% by weight, and preferably from 20 to 90% by weight, of at least one kind of a thermoplastic resin selected from a polyphenylene sulfide series resin, a polyamide series resin, a polyester series resin, a polycarbonate series resin, a vinyl chloride series resin, an olefinic resin, a polyacetal series resin, a polyarylate series resin, and a polyphenylene ether series resin;

A thermoplastic resin composition comprising

(A) from 5 to 90% by weight, and preferably from 10 to 80% by weight, of the graft polymer obtained 40 by graft polymerizing from 95 to 10% by weight of sum of the carboxyl group-containing vinyl monomer and the other vinylic monomer to from 5 to 90% by weight of the modified polyorganosiloxane (III),

(B) from 0 to 80% by weight, and preferably from 5 to 60% by weight, of the graft copolymer obtained by graft polymerizing from 95 to 10% by weight of the vinyl monomer excluding the carboxyl group-containing vinyl monomer to from 5 to 90% by weight of the polyorganosiloxane series polymer.

(C) from 0 to 80% by weight, and preferably from 3 to 70% by weight, of a styrene series resin, and

(D) from 10 to 90% by weight, and preferably from 20 to 80% by weight, of at least one kind of a thermoplastic resin selected from a polyamide series resin, a polyester series resin, a polycarbonate series resin, a vinyl chloride series resin, an olefin series resin, a polyacetal series resin, a polyphenylene sulfide series resin, a polyarylate series resin, and a polyphenylene ether series resin.

The styrene series resin (C) may be a composition composed of a resin obtained by polymerizing a monomer as a resin component in the presence of a rubbery polymer and a resin obtained by polymenzing a monomer as a resin component in the absence of the rubbery polymer.

Specific examples of the styrene series resin (C) are an acrylonitrile-butadiene rubber-styrene resin (ABS resin), an acrylonitrile-ethylenepropylene series rubber-styrene resin (AES resin), an acrylonitrilebutadiene-methyl methacrylate-styrene resin (ABSM resin), an acrylonitrile-styrene copolymer (AS resin), a methyl methacrylate-styrene copolymer (MS resin), high impact polystyrene (HIPS), an acrylonitrile-n-butyl acrylate rubber-styrene resin (AAS resin), etc.

For obtaining the thermoplastic resin composition of this invention described above, the above-

described components (Al. (B), etc., are mixed by, for example, a mixer and the mixture is melt-kneaded by an extruder at a temperature of from 20° C to 32° C and its formed into pellets. Furthermore, in a simple procedure, the above-described components can be directly melt-kneaded in a mold and molded into nellets.

The thermoplastic resin composition of this invention may, if necessary, contain an antioxidant such as 2.6-dit-butyl-4-methylp-henol. 2.1-methylcyclohexyly-4.6-dimethylphenol. 2.2-methylphenol. 2.1-methylcyclohexyly-4.6-dimethylphenol. 2.2-methylphenyl bioxphile, etc.: an ultravolet absorber such as p-butylphenyl salicate. 2.2-dihydroxy-4-methoxybenzophenone. 2-(2-hydroxy-4-m-octoxyphenyl)benzotriazole, etc.: a lubricant such as paraffin wax, stearic acid, hardened on, stearoamide, methylenebis-stearoamide, n-butyl stearate, ketone wax, octyl alcohol, hydroxystearic and Inglycenide, etc.: a flame retarder such as antimony oxide, chloroplatinic acid, alumnum hydroxide, zinc borate, tricresyl phosphate, trisdichlonopropyl) phosphate, chloronated paraffin, letrabromobutane, hexabromobenzene, letrabrombisphenol A, etc.: an antistatic agent such as stearoamide propyldimethyl-4-hydroxyethyl ammonum nitrate, etc.: a coloring inhibitor such as titanum oxide, carbon black, etc.: a filler such as calcium carbonate, day, silica, glass fibers, glass balls, carbon fibers, etc.: a coment, etc.

The pellets of the thermoplastic resin composition thus formed are molded by a conventional means such as compression molding, injection molding, etc.

As described above, the thermoplastic resin of this invention contains the graft copolymer formed by graft polymerizing a vinyl monomer to the modified polyorganosiloxane having bonded thereto a specific graft crosslinking agent, so that the graft polymerization of the vinyl monomer tends to occur as compared with the case of using a conventional polyorganosiloxane and the graft copolymer contained therein is formed with very high graft ratio and graft efficiency. Therefore, the thermoplastic resin of this invention containing the graft copolymer or the thermoplastic resin composition of this invention composed of the thermoplastic resin and other polymer resin is excellent in the balance of physical properties.

In particular, the thermoplastic resin or the thermoplastic resin composition has excellent cold resistance, weather resistance, slidability, abrasion resistance, impact resistance, moldability, weld strength and flame retardant. Of those properties, the slidability and abrasion reststance of the thermoplastic resin or the resin composition are higher than those of polyacetal and polyamide which are known as a sliding materials. Further, other properties of the resin or resin composition are equal or superior to those of an 35 ABS resin which is known to have balanced properties.

Since the thermoplastic resin or the thermoplastic resin composition of this invention has the excellent properties as described above, the resin or the resin composition can be applied to new fields such as sliding parts, parts for cold district, outdoor parts, etc., and the industrial significance thereof is very large.

This invention is described more practically by the following examples. In addition, all percents, parts.

35 ratios and the like are by weight unless otherwise indicated.

Various measurement items in the examples were obtained by the following manners.

That is, the graft ratio and the graft efficiency were determined by the following method.

A definite amount (x) of the graft polymerization product was added to acetone, followed by shaking by a shaker for 2 hours to dissolve the free copolymer, and the mixture was subjected to a centrifugal separation using a centrifugal separation at a rotation rate of 23,000 r.p.m. for 30 minutes to obtain insoluble matters. The insoluble product was dried for 1 hour at 120 °C using a vacuum dryer to provide an amount (y) of the insoluble product, and the graft ratio and the graft efficiency were calculated by the following equations.

Graft ratio = 
$$\frac{(y) - (x) \times (0)}{(x) \times (0)} \times 100(\$)$$
  
Graft efficiency =  $\frac{(y) - (x) \times (0)}{(x) - (x) \times (0)} \times 100(\$)$ 

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(Q): Content of component (III) in the graft polymerization product

Evaluations of the properties of the thermoplastic resin (composition) was made according to the following evaluation method (a) or (b).

Property	Unit	Test Method	Test Condition
Tensile Strength Bending Strength Bending Elasticity Break Elongation	kgf.cm² kgf.cm² kgf.cm²	ASTM-D638 ASTM-D790 ASTM-D790 ASTM-D638	Tensile rate 50 mm.min. Bending rate 15 mm.min. Bending rate 15 mm.min. Tensile rate 50 mm.min.
Izod Impact Strength (notched)	kgf * cm/cm	ASTM-D256	1 4", 23 °C 1 4", -30 °C
Heat Distortion Temperature Specific Gravity Combustibility Gloss Melt Flow Hate lockwell Hardness	°C	ASTM-D648 ASTM-D792 UL 94 ASTM-D523 JIS-K7210 ASTM-D785	Load 18.6 kgf cm <sup>2</sup> no annea 23° C.23° C - 45° 220° C 10 kg

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## Evaluation Method (b)

#### 25 Falling Weight Impact Strength:

An impact rod having a top curvature R of 1.2" was let fall from a height of 50 cm using an impact term and by Du Pont and the falling weight impact strength of each molded product having a thickness of 3.2 mm was measured. The unit was kgf cm.

## Sliding Character:

A friction rubbing test was made using a Suzuki type slide tester, and as a material to be rubbed with 35 the product, the same material as the product of steel (545C) was used. A test piece was a hollow cylindrical piece having an outer diameter of 25.6 mm and an inner diameter of 20.0 mm, and the form of the material to be rubbed with the test piece was the same as the test piece.

The kinetic friction coefficient was measured in an atmosphere at 23°C and 50% RH under a load of 5 kg and a running speed of 3.75 cm/sec.

The kinetic friction coefficient was calculated by the following equation.

$$\mu = \frac{3 \times F \times (r_2^2 - r_1^2)}{P \times (r_2^3 - r_1^3)}$$

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in the above equation. #represents a kinetic friction coefficient, F represents a force given to the load cell, P represents a load, F1 represents an arm length upto the load cell, r, represents an inner diameter, and represents an unter diameter.

The friction coefficient was measured in an almosphere of 23 °C and 50% RH; at a load of 5 kg, a running speed of 3.75 cm/sec.. and 12.600 rounds (running distance 0.24 km) in the case of using the same material as the sample as the material to be rubbed with the sample, and at a load of 10 kg, a running speed of 15 cm/sec, and 80,000 rounds (running distance 6 km) in the case of using steel as the material to be rubbed with the sample.

The friction coefficient was calculated by the following equation:

$$A = \frac{\Delta W}{P \times \ell \times \alpha}$$

wherein A represents a friction coefficient,  $\Delta W$  is a weight change of a sample, P is a load, t is a running distance, and  $\sigma$  is the density of a sample.

## Weather Resistance Test Method:

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After exposing each sample for 200 hours using a sun shine weather meter (Type WE-USN-HC, made by Toyo Rika K.K.) (exposure conditions: 63 °C, sometimes rain), Izod impact strength was measured.

#### Example 1

After mixing 1.5 parts of p-vinylphenylmethyldimethoxysilane and 98.5 parts of octamethylcyclotetrasiloxane, the mixture was added to 300 parts of distilled water having dissolved therein 2.0 parts of dodecylbenzenesulfonic acid and dispersed by emulsification by stirring for 3 minutes using a homo-

The mixture was placed in a separable flask equipped with a condenser, a nitrogen gas inlet and a streer, and heated to 90 °C for 6 hours with strring, followed by cooling for 24 hours at 5 °C to complete the condensation

The degree of condensation of the octamethylcyclotetrasiloxane in the modified polyorganosiloxane (III) obtained was 92 Re.

The modified polyorganosiloxane latex was neutralized to pH 7 with an aqueous sodium carbonate solution

35°arts of the modified polyorganosiloxane latex as a solids content was mixed with 0.5 part of sodium dodecy/benzenesulfonate and 140 parts of distilled water. the mixture was placed in a separable flask odupped with a dropping bottle, a condensor, a nitrogen gas inlet and a strere. After adding thereto 15.81 parts of styrene corresponding to 34% of the whole styrene amount, 6.29 parts of acrylonitrile corresponding to 34% of the whole scyrlonitrile amount, 0.2 part of sodium pyrophosphate, 0.25 part of grape sugar, 0.004 part of ferrous suitale and 0.074 part of cumene hydroperoxide, the temperature of the resultant mixture was raised to 70°C white flowing a nitrogen gas. After conducting the polymerization for 1 hour, a mixture was raised to 70°C white flowing a nitrogen gas. After conducting the polymerization for 1 hour, and the decomposed of 30.89 parts of remaining styrene, 1.221 parts of remaining arylonitrile, 1.084 parts of sodium dodecy/benzenesulfonate, 42 parts of distilled water, 0.12 part of cumene hydroperoxide, and 0.08 part of todecy/mercaptan was added thereto using the dropping bottle over a period of 3 numbers and 0.08 part of todecy/mercaptan was added thereto using the dropping bottle over a period of 3 numbers to the polymerization reaction was then further conducted for 1 hour and the reaction product was cooled.

The graft copolymer lake thus obtained was poured into warm water having dissolved therein 2 parts of calcium chloride dihydrate to conduct salting out coagulation and a thermoplastic resin containing the graft copolymer was separated. The thermoplastic resin was washed well with water and then drief for 16 hours at 80 °C to finish the nurrification.

57% of the resulting themoplastic resin powder was mixed with 43% of a copolymer (AS resin) obtained by emulsion polymerizing styrene and acrylonitrile at a monomer compounding ratio of 75:25 to provide a themoplastic resin composition. The thermoplastic resin composition was extrusion-molded using a twin screw extruder at a cylinder temperature of 230°C to provide pellets.

The thermoplastic resince composition obtained had excellent cold resistance, weather resistance, sliding property, impact resistance, and appearance. The evaluation results thereof are shown in Table 1 below.

# Examples 2 and 3 and Comparison Examples 1 and 2

By following the same procedure as in Example 1 except that the compounding ratio of the polyorganosiloxane (i) and the graft crosslinking agent (ii) was changed as shown in Table 1 below, the modified polyorganosiloxane latex and the thermoplastic resin containing the graft copolymer were pre-

pared. Using the thermoplastic resin, a thermoplastic resin composition was prepared by the same manner as in Example 1.

The evaluation results on these samples are also shown in Table 1.

In Comparison Example 1, the amount of the graft crosslinking agent was 0.1%. Therefore, the graft 5 ratio was reduced and a sufficient impact strength was not obtained. In Comparison Example 2, the amount of the graft crosslinking agent was 15%. Therefore, the molecular weight of the vinyl monomer grafted to the modified polyorganosiloxane and the vinyl monomer not grafted was reduced and a sufficient impact strength was not obtained.

## Comparison Examples 3 to 5

By following the same procedure as in Example 1 except that 1.5 parts of vinylmethyldimethoxysilane. 15 y-mercaptopropylmethyldimethoxysilane or y-methacryloxypropylmethyldimethoxysilane was used in place of p-vinylphenylmethyldimethoxysilane at the preparation of the modified polyorganosiloxane (III), a modified polyorganosiloxane latex, a thermoplastic resin containing a graft copolymer (V) and a thermoplastic resin composition were prepared.

Results of the evaluation on these thermoplastic resins in the same manner as in Example 1 are shown 20 in Table 1 below.

In Comparison Examples 3 and 4, a high graft ratio was not obtained and a sufficient impact strength was not obtained

In Comparison Example 5, almost the same impact strength as the product in Example 1 was obtained, but the appearance was poor and sufficient gloss was not obtained.

#### Examples 4 to 6

By following the same procedure as Example 1 except that the vinyl monomer (IV) grafted to the modified polyorganosiloxane (III) was changed as shown in Table 1 below and the resin blended with a thermoplastic resin obtained was changed as shown in Table 1, each thermoplastic resin composition was prepared. The evaluation results of these resin compositions are shown in Table 1.

As is clear from the results shown in Table 1, the graft copolymers (V) had a high graft ratio and 35 thermoplastic resin compositions having a high impact strength and good appearance were obtained.

#### Examples 7 and 8

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By following the same procedure as Example 1 except that the graft crosslinking agent was changed as shown in Table 1, each thermoplastic resin and thermoplastic resin composition were prepared and evaluated. The results are shown in Table 1.

As is clear from Table 1, the graft copolymers (V) had a high graft ratio and thermoplastic resin compositions having a high impact strength and a good appearance were obtained.

			5	.0	20	15	10	
		TAE	TABLE 1					i
Blending Composition (part) at Preparation of Component (111)	Example 1	Example 2	Example 2 Example 7	Example 4	Example 5	Example 6	Example 1	Example
Component (I) Octamethylcyclotetrasiloxane								
Component (II)	5.86	99.25	97.0	99,5	5, 86			
2-(p-vinylph amenydamethoxysilane 3-(p-vinylbenzoyloxy)prophyldimethoxysilane	1.5	0.75	3.0	:			90.5	98.5
Vinylacthoxyailane		٠.	, ,		1.5	2.5	1 -	
Y-Methacryloxypropylmethoxysilane				,				1.5
Blending Composition (part) at Preparation of Component (IV)			ı			. 1		
Component (111) Latex (solid conversion) Styrene Actylonitile Worly! Hethacrjate	35 46.5 10.5	35 46.5 10.5	35 18.5	8.80 i i	8. 1.8	35 45.5 2.9.5	35 186.5 18.5	35 18.5
Composition(%) of Thermoplastic Resin Composition		٠						
The moplastic Resin of the Invention Resident of 1975(1) The Polymethylaed Resident of 1975(1) The Stytems Resident of 1975(1) The Polymethylaed Resident of 1975(1) The 1975(1) The Polymethylaed Resident of 1975(1) The Polymethylaed Resident of 1975(1) The Polymethylaed Resident of 1975(1) The Pol	2 C	68 1 1 1 1	483	6.6	133.1	57	55,	43
Evaluation								
Graft ratio(%) of Component (V) Graft Efficiency (%)of Component (V)	101	51.	117	112	137	125 67	34	53
Property of Thermoplastic Resin Composition:								
Izod Impact Strength (23°C, kgf.cm/cm)	32.3	23.0	31.5	23.1	20.1	21.1	32.2	32.1
Gloss(%)	85	10	63	11	7.8	7.1	18	82

5		Comparison Example 5		98.5		1.5		35 186.5 5		57 43 		100 53		25.0	*
10		Comparison Exemple 4		98.5				335 18.55		43 1		9		5.7	9
·5 20		Comparison Example 3		98.5		8.1.1		38. 186. 18.5		437		25		4.0	
25	ont'd)	Comparison Example 2		9.2	15			24.05 20.05 20.05		43. 1		130		9.0	. 0.61
30	TABLE 1 (cont'd)	Comparison Example 1		6.66	1.0	ri.		35 1.66.5 1.55.55		437		2.5 8		3.0	31.0
35															
40		t Preparation			rsilane Idimethoxysilane methyldimethoxy-	hoxysilane imethoxysilane	it Preparation	conversion)	ic Resin	nvention		(V) ment (V)	in Composition:	kgf.cm/cm)	
45		art) a	loxane		methy ropyl	ilane Idimet ethyld	art) e	(solid	oplast	the I 5(8)] Resin		Compo	ic Res	23°C,	
50 55		Blending Composition (part) at Preparation of Component (III)	Component (1) Octamethylcyclotetrasiloxane	Component (II)	2-(p-vinylphenyl)ethylmethoxysilane 3-(p-vinylbenzoyloxy)propylmethyldimethoxysilane silane	Vinylmethyldimethoxysilane Y-Mercaptopropylmethyldimethoxysilane Y-Methacryloxypropylmethyldimethoxysilane	Blending Composition (part) at Preparation of Component (IV)	Component (III) Latex (solid conversion) Sylvene Acrylenitile Acrylenitile Hethyl Hethacrylate	Composition(%) of Thermoplastic Resin Composition	Thermoplastic Resin of the Invention Resin (St. 1875[4]) Polystycen Resin Polystycen Resin Stylesthylmethacytate Resin Styrene(MHA Copolymer Resin STYAMA = 70/30[4])	Evaluation	Graft ratio(%) of Component (V) Graft Efficiency (%)of Component (V)	Property of Thermoplastic Resin Composition:	Izod Impact Strength (23°C, kgf·cm/cm)	Gloss(%)
		P of	ຽັ	3 -		-	E o	Z>NC	ပိပိ	- <r< td=""><td>짋</td><td>00</td><td>P</td><td>-</td><td>9</td></r<>	짋	00	P	-	9

For examining the differences between the product in Example 1 and the product in Companson Example 5, the following companson tests were prepared.

- The change of the graft ratio in the case of molding each product under the temperature condition
   shown in Table 2 is shown in Table 2 below.
  - In Example 1, the graft ratio at the end of the polymerization is almost the same as the graft ratio after molding at 280 °C, which shows the graft point is thermally stable.
  - On the other hand, in Companson Example 5, the graft ratio at the end of the polymerization is reduced when the product is molified at 230°C, which shows that the graft point is thermally unstable.

TARLE 2

*5		End of Polymerization	230 `	260 °C	280°C	300°C
	Graft Ratio in Example 1	101	100	100	99	92
	Graft Ratio in Comparison Example 5	100	96	85	76	66

in Table 2, the temperature is molding temperature.

- (2) The falling weight impact strength in the case of molding each product under the temperature condition shown in Table 3 is shown in Table 3 below.
- It is considered that the reduction of the graft ratio clarified in above Test (1) gives an influence on the falling weight impact strength.
  - In Example 1, the reduction of the strength is small in molding at 280 °C, while in Comparison Example 5, the strength is greatly reduced in molding at 280 °C or higher as compared with molding at 260 °C.

TABLE 3

	260 ° C	280 °C	300 °C
	Molding	Molding	Molding
Falling Weight Impact Strength in Example 1 Falling Weight Impact Strength in Comparison Example 5	330	320	250
	325	260	100

(3) For observing the retention heat stability in a molding machine, the reduction of gloss from the 1st shot to the 5th shot and the recovering property thereof in continuous molding in the case of conducting injection modding by repeating at a cycle of 60 seconds and in the case of once stopping the molding machine and after retaining the product therein for 15 minutes, conducting injection molding at a cycle of 60 seconds were examinad.

The evaluation is made at a molding temperature of 260 °C and 280 °C.

The results are showing in Table 4 below.

In Example 1, the change of gloss is small and the recovering property thereof is good. On the other hand, in Comparison Example 5, the change of gloss is observed at a molding temperature of 260°C and when molding is conducted at 280°C, the gloss is extremely reduced.

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TABLE 4

				Gloss(°	6)		
	Molding temperature	Continuous molding	1st shot	2nd shot	3rd shot	4th shot	5th shot
1	(, C)						
Example 1	260 280	84.0 83.5	83.5 81.0	83.0 80.5	81.0 76.5	82.5 79.5	84.0 81.5
Comparison Example 5	260 280	84.0 82.0	80.0 66.5	80.0 63.0	75.0 55.0	77.0 66.0	81.0 76.0

After retention of 5 minutes

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## Example 9 and Companson Example 6

In Example 9, a thermoplastic resin composition composed of 51% of the thermoplastic resin obtained in Example 1 and 49% of the AS resin as used in Example 1 was molded by the same method as in Example 1.

The content of the modified polyorganosiloxane (III) in the thermoplastic resin composition of Example 9 was 17.85%.

On the other hand, the resin in Comparison Example 6 was an ABS resin (JSR ABS12, trade name; made by Japan Synthetic Rubber Co., Ltd.) and the content of butadiene rubber in the resin was 17.8%.

On these resins, the properties were compared according to the evaluation methods (a) and (b). The 30 results obtained are shown in Table 5 below.

It can be seen that the thermoplastic resin of this invention has particularly excellent cold resistance. weather resistance and sliding property as compared with the commercially available ABS resin, and also other properties of the thermoplastic resin of this invention are same as or higher than those of the ABS resin.

Furthermore, 100 parts of each of the resins in Example 9 and Comparison Example 6 was blended with 10 parts of tetrabromobisphenol A, 3 parts of antimony trioxide and 0.0001 part of chloroplatinic acid, and a flame retardant test was made on each blend according to UL-94.

The results are shown in Table 5 below.

It can be seen that the thermoplastic resin of this invention shows an excellent flame retardance by the 40 addition of a small amount of flame retarder, which gives the merits of inhibiting the reduction of general properties, reducing the toxicity, and also reducing the volume cost, and hence an excellent flame retardant resin is obtained.

## TABLE 5

5		Example 9	Comparison Example 6
	Izod Impact Strength (23°C)	32.3	37.2
	Falling Weight Impact Strength	338	430
10	Heat Distortion Temperature	95	91
	Rockwell Hardness	102.1	106.0
:5	Melt Flow Rate	4.2	13.5
	Tensile Strength	430	490
	Break Elongation	34	36
20	Bending Strength	720	790
	Bending Elasticity	24,500	26,700
25	Gloss	81	78
	Izod Impact Strength (-30°C)	14.1	11.5

## TABLE 5 (cont'd)

5		Example 9	Comparison Example 6
3	Izod Impact Strength After Weather Resistance Test	28.8	25.1
10	Kinetic Friction Coefficient with the Same Material as the Test Sample	0.09	0.29
15	Kinetic Friction Coefficient with Steel	0.09	0.46
	Wear Coefficient with the Same Material as the Test Sample	16×10 <sup>-3</sup>	498×10 <sup>-3</sup>
20	Wear Coefficient with Steel	19×10 <sup>-3</sup>	621×10 <sup>-3</sup>
	Combustibility	HB passed	HB passed
	Specific Gravity	1.055	1.021
25	Flame Retardability		
	· HB Test	passed	passed
30	V-2 Test	passed	Failed
	V-1 Test	passed	Failed
	V-0 Test	passed	Failed

## Examples 10 and 11

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In these examples, the thermoplastic resins were molded without blending with other resin, and evaluated. The polymerization condition, the molding condition and the evaluation condition were same as those in Example 1. The results are shown in Table 6.

TABLE 6

Blending Composition (part) at Preparation of Component (III)	Example 10	Example 11
Octamethylcyclotetrasiloxane p-Vinylphenylmethyldimethoxysilane	98.5 1.5	98.5 1.5
Blending Composition (part) at Preparation of Component (IV)		
Latex of Component (III) (solids content) Styrene Acrylonitnle Methyl Methacrylate	20 60 20	20 - - 100
Evaluation		
Graft ratio (%) of Component (IV) Graft Efficiency (%) of Component (IV)	140 35	120 30
Property of Thermoplastic Resin:		
Izod Impact Strength (23 °C, kgf cm cm) Gloss	28.0 80	19.6 82

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#### Example 12

After mixing 20 parts (solids content) of the modified polyorganoslioxane latex obtained in Example 1, 0.5 part of sodium dodecylbenzernesulfonate and 140 parts of distilled water, the mixinger was placed in a separable flask equipped with a condenser, a nitrogen gas inlet and a stirrer. After adding thereto 34 parts of styrene, 46 parts of N-phenylmaleimide, 0.1 part of sodium ethylenediaminetetracetate, 0.003 part of ferrous suitate and 0.1 part of diisopropry) benzene hydroperoxide, the emperature of the resultant mixture was raised to 70 C while flowing a nitrogen gas, followed by conducting polymerization for 1 hour. Thereafter, the reaction mixture was cooled.

2 Parts of the graft copolymer latex thus obtained was added to pressurized warm water at 130 °C having dissolved therein calcium chloride dihydrate to conduct salting out and coagulation, and a thermoplastic risin containing the graft copolymer was separated.

The thermoplastic resin was washed well with water and then dried at 80°C for 16 hours for the punification, and the product was molded into pellets by an extruder having a cylinder temperature of 280°C.

Store graft ratio of the thermoplastic resin obtained was 120%, and the thermoplastic resin had a heat distorion temperature of 160 °C, an Izod impact strength of 8 kgf cm/cm and a melt flow rate at 280 °C of 10 gr/10 min.

#### Example 13

After mixing 1.5 parts of p-vinylphenyldimethoxysiane and 98.5 parts of octamethylcyclotetrasiloxane, the mixture was added to 300 parts of distilled water having dissolved therein 2.0 parts of dodecylbenzene sulforate. After roughly emulstyring the mixture by stirring for 5 minutes using a homomixer, 3.0 parts of a 50% methanol solution of 4/3-triethoxysilylpropoxyl-2-hydroxybenzophenone was added hereto, followed by stirring for 5 minutes. The mixture was then dispersed by emulsification using a press homogenizer.

The mixture was placed in a separable flask equipped with a condenser, a nitrogen gas inlet and a stirrer, heated to 85°C for 5 hours with stirring, and then allowed to stand for 12 hours at room temperature

to complete the polycondersation.

The product was neutralized to pH 7 with a 10% aqueous solution of sodium carbonate. The degree of The product was neutralized to pH 7 with a 10% aqueous solution of somethylocytobersalioxane in the modified polyorganosiloxane was 92.3%, the average sondensation of cotamethylocytobersalioxane was 92.3% the content of the pdegree of polymerization of the product was 5.200 as a silicon atom number, the content of the UV absorbing virilybhenyl group in the organic groups in the polymer was 0.27%, and the content of the UV absorbing group was 0.13%.

Using the modified polyorgenoslovane latex thus obtained, styrene and acrylonitrile were graft polymerUsing the modified polyorgenoslovane latex thus obtained, styrene and acrylonitrile were graft polymerto lates the

A thermoplastic resin composition was prepared by blending the polysiloxane-modified thermoplastic resin obtained by the above method with other organic polymer.

That is, the resin composition was prepared by mixing 57% of the powder of the polyorganosioxanemodified thermoplastic resin obtained and 43% of a copolymer (AS resin) obtained by emulsion polymodified thermoplastic resin obtained and 43% of a copolymer (AS resin) obtained by emulsion polyin metizing stryene and acrylontine in a monomer blending ratio of 75 : 25 by weight ratio. The combination of the above components is shown in Table 7 below.

The thermoplastic resin composition was extruded into pellets using a twin screw extruder at a cylinder temperature of 230°C,

On the thermoplastic resin composition thus obtained, the impact resistance, sliding property, weather On the thermoplastic resin composition.

resistance and gloss were measured. In addition, the weather resistance test was conducted as follows. That is, after exposing the sample for in addition, the weather resistance test was conducted as follows. That is, after exposing the sample for in addition, the weather meter (Type WE-USN-HC, made by Toyo Rika K.K.), 1500 hours (83° C, no rain) using a sun shine weather meter (Type WE-USN-HC, made by Toyo Rika K.K.), and the discolored degree Lab (L. brightness, at redness, b: yellowness) was measured by a colorimeter of Type AU-SCH-2, made by Suga Shikenki K.K., and the tone changed value E was calculated by the following returning.

$$E = \sqrt{(L_1 - L_2)^2 + (a_1 - a_2)^2 + (b_1 - b_2)^2}$$

wherein L<sub>1</sub>, a<sub>1</sub> and b<sub>1</sub> are tone before the weather resistance test, and L<sub>2</sub>, a<sub>2</sub> and b<sub>2</sub> are tone after the weather resistance less:

In the above equation, the lower value of E shows that the change of color is small, that is, the resin composition has excellent weather resistance.

#### Example 14

By following the same procedure as in Example 13 except that 1.5 parts of -methacryloxypropylmethyldimethoxysitane was used in place of p-iniylphenylmethyldimethoxysiane and 3.0 parts of a 50% methanol solution of 4-0-methyldiethoxysitylpropoxyy-2-hydroxybenzophenone was used in place of the 50% methanol solution of 4-0-methyldiethoxysitylpropoxyy-2-hydroxybenzophenone, a polyorganosiloxane-modi-50% methanol solution of 4-0-d-retriboxysitylpropoxyy-2-hydroxybenzophenone, a

s fied thermoplastic resin was prepared.

The average degree of polymerization of polyorganosiloxane was 3,900 as a silicon atom number, the content of the reactive unsaturated group in the organic groups of the polymer was 0.24%, and the content of the UV absorbing group was 0.15%.

Furthermore, a thermoplastic resin composition was prepared by blending the polyorganosiloxanemodified thermoplastic resin and the AS resin under the same condition as in Example 13. The combination
of the above components is shown in Table 7 below.

On the thermoplastic resin composition thus obtained, the impact resistance, sliding property, weather resistance and gloss were measured in the same manner as in Example 13.

The results are shown in Table 7 below.

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#### Example 15

By following the same procedure as in Example 13 except that 1.0 part of methylvnryldimethoxysilane was used in place of p-vnrylphenylmethyldimethoxysilane and 2.0 parts of laurylsulfunc acid was used in place of dodecylbenzenesulfonc acid. a polyorganosiloxane-modified thermoplastic resin was prepared.

The average degree of polymerization of the polyorganosiloxane was 5,400 as a silicon atom number. The average degree of polymerization of the organic groups in the polymer was 0,28%, and the content of the reactive unsaturated group in the organic groups in the polymer was 0,28%, and the content of the UV absorbing orgoup was 0,14%.

Furthermore, a thermoplastic resin composition was prepared by blending the polyorganosiloxanemodified thermoplastic resin and the AS resin under the same condition as in Example 13.

On the thermoplastic resin are set of the composition obtained, the impact resistance, sliding property, weather resistance and gloss were measured. The results are shown in Table 7 below.

## Examples 16 to 18 and Reference Example 1

The same procedure as Example 13 was followed using the components shown in Table 7 below. The evaluation results of these products are shown in Table 7.

It can be seen from the results shown in Table 7 that the thermoplastic resin composition containing the graft copolymer having the UV absorbing group has excellent wealher resistance.

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5		Reference Example		98.5	1.5			35 46.5 18.5
10		Example 18		98.5	1.5	3.0		35 45.5 19.5
15		Example 17		98.5	1.5	3.0		35
		Example 16		98.5	1.5	9.0		35
20		Example 15		9.8.5	1.5	3.0		35 46.5 18.5
25	7	Example 14		98.5	, 1.5	3.0		35 46.5 18.5
30	TABLE 7	Example 13		90.5	1.5	3.0		35 46.5 18.5
35					lane Lhoxysilane	508 Hethanol Solution of 4-(3-triethoxy- sillypropoyy)2-hydroxybensoplenone 508 Methanol Solution of 4-(3-methyldiethoxy- sillypropoxy)-2-hydroxybenzophenone		lymer nt (A)
40				iloxane	p-Vinylphenylmethyldimethoxysilane Y-Wethacryloxypropylmethyldimethoxysilane Methylvinyldimethoxysilane	500 Methanol Solution of 4-(3-triethoxy- silyhytoxyy)-2-Widkoxybensophenone 500 Methanol Solution of 4-(3-methyldieth silyhpropoxy)-2-hydroxybenzophenone		Blending Composition for Graft Copolymer Polyorganosiloxane of Component (A) Styrene Accylonitrile Mathyl Methacrylate
45		Polyorganosiloxane-Modified Thermoplastic Resin:	art)	<ul><li>(a) Octamethylcyclotetrsiloxane</li></ul>	p-Vinylphenylmethyldimethov Y-Methacryloxypropylmethyld Methylvinyldimethoxysilane	nol Solutic oxy)-2-hyd nol Solutic oxy)-2-hyd	part)	ng Composition for Polyorganosiloxane Styrene Acrylonitrile Wethyl Methacrylate
50		Polyorganosiloxane-Ho Thermoplastic Resin:	Component (A) (part)	Octamethy!	p-Vinylpho y-Methacrj Methylviny	50% Methau silylprop 50% Methau silylprop	Component (B) (part)	ling Composition Polyorganosilo Styrene Acrylonitrile Methyl Methaci
55		Polyorga Thermop)	Сомрол	(a)	(a)	(c)	Compor	ВІепс

				•		10	5
TAI	1 7 (	TABLE 7 (cont'd)					
Thermoplastic Resin Composition and Evaluation Thereof:	Example 13.	Example 14	Example 15	Example 16	Example 17	Example 18	Reference Example
Composition (8)							
Blend Resin:							
Polyorganosiloxane-modified Thermoplastic Resin	57	75	5.1	53	57	57	5.7
NS Resin Polystyrene Resin Polymethyl Methacrylate Resin Styrene/MMA Copolymer Resin		£	£3	£	43	64	43
Evaluation:							
Graft Ratio(a) of Graft Copolymer	101	99	53	100	9.1	100	102
Graft Efficiency(8) of Graft Copolymer	54	53	29	54	52	54	55
Trad Impact Strength (23°C)	32	33	20	19	16	23	32
_	14	13	6	6	89	70	14
Kinetic Priction Coefficient with the same	0.09	0.09	0.09	0.09	0.09	0.0	0.09
Material as the Sample Relative Worn Amount with the same Material	16	16	16	16	16	16	16
<pre>(×10<sup>-2</sup>) Izod Impact Strength After Weather</pre>	59	28	1.1	16	13	20	29
Resistance Test (kgf.cm/cm) Appearance Change After Weather Resistance	3.1	3.2	3.1	4.5	2.6	1.5	10.8
Test ( $\Delta E$ ) Gloss (%)	8.5	84	7.2	83	83	8.5	9.2

## Examples 19 to 34 and Comparison Examples 7 to 9

By following the same procedure as Example 13 using the components shown in Table 8 below, each of polyorganosiloxane series polymers R-1, R-2, and R-3 were prepared.

TABLE 8

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	R-1	R-2	R-3
p-Vinylphenylmethyldimethoxysilane (part)	1.5	0.05	55
Octamethylcyclotetrasiloxane (part)	98.5	99.95	45
Polymerization Conversion (%)	98.0	97.5	98.5

By following the same procedure as in Example 13 according to the compositions shown in Table 9 below using the polyorganosiloxane series polymers prepared in the above step, graft copolymers G-1 to G-8 were prepared.

TABLE 9	(Cont 'd	4					
G-1	C-5	6-3		G-5	9-9	G-7	G-8
					;	;	;
60 35	35	35	09 -	ا و0	30 00	35	40
1 1	1 1	1 1	32	35	1 1	1 1	1 1
1 1	1 1	1 1	1 1	1 1	ın ı	ıvı	1 1
'n.	ហ	ហ	ហ	v		1	1
30	S	35	28	25	30	29	31
	G-1 G-1 60 35 - - 5	#BBLE 9 (Contrid	### 9 (Cont.d)    G-1   G-2   G-3     G-2   G-3     G-3   G-3     G-1   G-3     G-1   G-3     G-2   G-3     G-2   G-3     G-3   G-3     G-3	60 60 35 5 5 35	G-3 G-4 35 35 17 17 17 17 17 18 18 18 18 18 18 18 18 18 18	60 60 60 60 60 60 60 60 60 60 60 60 60 6	60 60 60 60 60 60 60 60 60 60 60 60 60 6

Each composition shown in Table 9 below was melt-kneaded by an extruder having an inner diameter of 40 mm at a lemograture range of from 280 to 320 °C to provide pellets.

The pellets were moided using a 5oz njection molding machine (1S-80A, made by Toshiba Corporation) at the temperature range of from 280 to 320°C to provide each test piece, and the properties of each sample were evaluated. The results are shown in Table 9 below.

In addition, the heat distortion temperature was measured under conditions of a load of 18.6 kg/cm<sup>2</sup> and no anneal according to ASTM D648.

The chemical resistance was evaluated by applying a definite strain of 1% in strain ratio to each test picce (18".12", 5"), coating a brake fluid composed of dicctyl phthalate (DOP) on the strained portion, and after allowing to stand the test piece at 23 °C and measuring the time until the test piece was ruptured, the time was employed as the index of the chemical resistance.

In the above evaluation, o shows the case of causing neither crack nor rupture over 100 hours.

The weld strength retentivity was obtained as follows.

That is, the tensile strength (Tw) of the test piece was measured on the molded test piece using a mold appearing a weld line at the center of ASTM No. I Dumbbell, the tensile strength (To) of the test piece was measured on the test piece molded using a mold of not giving the weld line, and the weld strength retensitivity of Tw To x (100% was determined.

The molding property was evaluated by the presence or absence of the occurrence of gellation or the 
ropture of moldings in the step of granulating the product by an extruding machine and obtaining the 
moldings by a molding machine.

The appearance of the moldings was evaluated by the presence or absence of the occurrence of a flow mark silver on the surface of the moldings.

In addition, in the examples and the comparison examples, the following materials were used as the composition (A).

PPS (polyphenylene sulfide series resin): Toprene T-4, made by Toprene K.K.

Nylon 6: Amilan CM 1017, made by Toray Industries, Co., Ltd.

Nylon 4.6: Stanyl KS300, made by DSM Co. PBT (polybutylene terephthalate): Duranex XD499 made by Polyplastic Co.

PC (polycarbonate): A-2200, made by Idemitsu Sekiyu-Kagaku K.K.)

PVC (polyvinyl chloride): A-2200. made by Toagosei Chemical Industry Co., Ltd.)

MPP (maleic anhydride-modified polypropylene), MODIC P-10B, made by Mitsubishi Peterochemical Comananu I id

POM (polyoxymethylene): Duracon M90, made by Polyplastic K.K.

Polyarylate: U Polymer U-8000, made by Unitika Ltd.

PPE (polyphenylene ether) was prepared by the following method.

## (1) Production of PPE-1 (polyphenylene ether):

Polymerization reaction of 2.6-xylenol was carried out in a toluene solution while blowing oxygen into the solution at 30°C using cupric bromide and di-n-butyleneamine as a catalyst and the polymerized product was purified to provide a PPE-1.

[7] of the polymer PPE-1 measured at 30 °C using chloroform as solvent was 0.40 dl.g.

## (2) Production of PPE-2 (maleic anhydride modified polyphenylene ether):

After dry-blending 100 parts by weight of the polymer PPE-1 with 2 parts of maleic anhydride and 1 part of 2-dimethyl-2.5-dift-butylenoperoxy)hexane at room temperature, the blend was melt-kneaded using a same direction rotary system twin screw extruder equipped with vent at a cylinder temperature of 300° C and a screw rotation number of 150° r.p.m. to form pellets of the polymer, i.e., maleic anhydride-modified polybrieneylene ether (PPE-2)

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TABLE 10-A	Example Example Example Composition: 20 Composition:	01 01	30	08			1	Amount (%) of Polyorganosiloxane in Composition 18 18	Amount (%) of Epoxy Group-Having Vinyl Monomor in 1.5 Composition	lzod Impact Strength (23°C) (kgf.cm/cm) 30 5 HFR (g/10 min., 300°C) 36 34	00 00 138 139 130 130 130 130 130 130 130 130 130 130	.09	poob
	le Example	70	. 1	. !	30	11	1	18	1.5	10 37	0082	103 100 0.09 0.09 .6×10 <sup>-3</sup> 16×10 <sup>-3</sup> 8	poob poob
	Example 22	70			1 05	311	ı	18	1.5	37	೦೦ಇ೯	0.09 16×10 <sup>-3</sup> 13	poob
	Example 23	7.0		, , ,		30	,	18	1.5	10 36	0082	7	poob
	Example 24	94	30	, , ,			30	18	1.5	2°e	೦೦೭೭	0.09 16×10 <sup>-3</sup> 6	pood

TABLE 10-A (cont'd)

5 10 5	Example Exampl	07 07 07 07 07 07 07 07 07 07 07 07 07 0	30 30 30 30 30	Lyorganosiloxane in Composition 18 18 18 18	Amount (1) of Epoxy Group-Having Vinyl Monomer in 1.5 1.5 1.5 1.5 1.5 Composition	Aluation Result:  Tand Impact Strength (23°C) (kgf.cm/cm)  Tand Impact Strength (23°C) (kgf.cm/cm)  Tand Indiana (210°m)  Tand India
	Blending Composition (Part) of Composition: Composition (Part) of Composition: Volum 6	PBT 4, 6 PC PVC PVC PQC PQC PQC PQC PQC PQC PQC PQC PQC PQ	Component (B) G-1	Amount (%) of Polyorganosiloxane in Composition	Amount (%) of Epoxy Group-Having Composition	Pvaluation Result: Isod Impact Strength (23°C) (Agf-cm/cm) Cham (40'10 min   Framp.) Cham (40'10 min   Framp.) Rese Puid Resistance: Rese Puid Strength Resentivity (8), Chamber of Strength Resentivity (8), Chamber of Strength Coefficient (vith same Mantise of Strength Agrand (4), Chamber (4

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Екатр је		30	10	1.5	30 (260) 32 (260) 0 (81) 10.09 16.09 16.09 2900d 900d
Examplo 33		30	18	1.5	23 34 (260) 0 78 182 100 16,09 22 200d 900d
Example 32	1111115	30	18	1.5	25 (300) 0 (300) 101 101,09 16110-3 2410-3
Example 31		30	1.8	1.5	18 (200) 0 O O O O O O O O O O O O O O O O O O O
Example 30	11111011111	30	1.8	1.5	24 (220) O O O B B B B B B B B B B B B B B B B B
Blending Composition (part) of Composition: Nichont (A) Nichont (A)	PET	Component (B)	Amount (8) of Polyorganosiloxane in Composition	Amount (%) of Epoxy Group-Having Vinyl Monomer in Composition	Evaluation Result:  130d Impact Strength (23°C) (Agf.cm/cm)  FOR (45/0 min.) (Temp.)  FOR (45/0 min.) (Temp.) (Temp.) (Temp.)  FOR (45/0 min.) (Temp.) (Temp.) (Temp.)  FOR (45/0 min.) (Temp.) (Temp.) (Temp.) (Temp.)  FOR (45/0 min.) (Temp.)

It can be seen from the results shown in Table 10 that in Examples 19 to 34, thermoplastic resin compositions desired in this invention are obtained. On the other hand, in Companson Examples 7 to 9, the desired effects of this invention cannot be obtained. That is, in Companson Examples 7 and 8, the amount of the graft crosslinking agent in each polyogranosiloxans series copolymer is outside the range defined in this invention and as a result, each product is inferior in Izod impact strength.

Reference Example 2 is an example of a composition outside the range of this invention, which was prepared using a graft copolymer without using an epoxy group-containing monomer. The impact strength, gloss, weld strength retentivity, moldability and the sppearance of the molding are inferior. Reference Example 3 is an example wherein the graft copolymer is less than the range of this invention and the product is inferior in impact resistance and weld strength retentivity. Reference Example 4 is an example wherein the graft copolymer is over the range defined in this invention and the product is inferior in heat distortion temperature.

Comparison Example 9 shows the properties of PPS (polyphenylene sulfide series resin).

#### Examples 35 to 60

By following the same procedure as in Example 1 using the components shown in Table 11 below, a polyorganosiloxane series copolymer R-1, R-2 and R-3 were obtained.

#### TABLE 11

	R-1	R-2	R-3
p-Vinylphenylmethyldimethoxysilane (part)	1.5	0.05	55
Octamethylcyclotetrasiloxane (part)	98.5	99.95	45
Polymerization Conversion (%)	98.0	97.5	98.5

By following the same procedure as in Example 13 using the above polyorganosiloxane series polymer according to the compositions shown in Tables 12 and 13 below, graft copolymers A-1 to A-8, B-1 to B-8 and C-1 to C-3 shown in Table 12 and Table 13 were obtained.

TABLE 12

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Polymer	A-1	A-2	A-3	A-4
Batch Blending Composition (part)				
Polyorganosiloxane Senes Polymer				
R-1 Styrene Acrylonutrile o-Methylstyrene Methacrylic acid Acrylic acide	40 14.3 4.7 - 1	40 11.3 4.3 3.3 1	40 14.3 4.7 - 1	40 19.0 - - 1
Blending Composition (part)				
Styrene Acrylonutrile Melthytstyrene Methacrylic acid Acrylic acid Polymerization (**)	28.7 9.3 - 2 - 98.5	22.7 8.7 6.7 2 - 98.3	28.7 9.3 - - 2 98.5	38.0 - - 2 - 98.9
Polymer Composition (%)	,			
Polyorganosiloxane Senes Polymer Styrene Acrylonitrile a-Methylstyrene Methacrylic acrd Acrylic acid Graft ratio (%)	40 43 14 - 3 - 95	40 34 13 10 3	40 43 14 - 3 94	40 57 - 3 - 98
Limiting Viscosity [n] (dl-g)	0.50	0.48	0.51	0.50

TABLE 13

	Polymer	B-1	B-2	B-3	C-1	C-2	C-3
5	Batch Blending Composition (part)						
	Polyorganosiloxane Series Polymer						
	R-1	40		40	-	-	-
	R-2		٠ .			•	•
10	R-3	-	40		-	•	-
	Styrene	15	15	12.5	75	67.5	100
	Acrylonitrile	5	5	4.2	25	22.5	-
	α-Methylstyrene			3.3		10	•
	Methacrylic acid	i -	٠ .			-	•
*5	Acrylic acid			·	•	-	<u> </u>
	Blending Composition (part)						
	Styrene	30	30	25		-	-
	Acrylonitrile	10	10	8.3	- 1	-	-
20	a-Methylstyrene	١ ٠		6.7		-	•
	Methacrylic acid	-	-		- '	-	•
	Acrylic acid	-					•
	Polymerization (%)	98.7	98.8	98.4	99.5	99.8	99.7
25	Polymer Composition (%)						
	Polyorganosiloxane Series Polymer	40	40	40	-	-	-
	Styrene	45	45	37.5	75	67.5	100
	Acrylonitrile	15	15	12.5	25	22.5	-
	α-Methylstyrene		-	10	-	10	
30	Methacrylic acid		-	-			
- 1	Gralt ratio (%)	96	102	88	-	-	-
- 1	Limiting Viscosity [7] (dl/g)	0.51	0.21	0.47	0.60	0.55	0.51

The components (A) to (D) shown in Table 14 below were melt-kneaded by an extruder having an inner diameter of 40 mm at the temperature range of from 250 to 320 °C to provide pellets.

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The pellets were moded using a 5oz injection molding machine (IS-80A, made by Toshiba Corporation at a molding temperature of from 260 to 300°C to provide each test piece and the properties of the test piece were evaluated. The results are shown in Table 14 below.

In addition, in the above examples and comparison examples, the component (D) used was the same as in Examples 19 to 34 described above.

As is clear from Table 14, in Examples 35 to 60, the desired thermoplastic resin compositions of this invention are obtained.

10		Example Example 20			10 70	10		10 10			- 10		50 50 50									16 16 16	0.3 0.3 0.9		1) 15 20 10 9 12 45 (240) 41 (240) 43 (240)	00	986	0-3 16	
·		Example 37		70	,		a			1			80								,	7.6	0.3		10 6 33 (240)	၀၁	85 78	0.09 16×10 <sup>-3</sup> 10	ł
20		Example 36		10	•		0	•		30			20	. ,						,	ı	20	0.3		20 12 ) 20 (240)	00	90	0.09 16×10 <sup>-3</sup> 19	1
25	FABLE 14	Example 35		10	٠,		30	. ,		9 '			20	٠,							•	16	0.3		18 11 36 (240)	00	8 8	0.09 16×10-3 17	;
. 30																		•				9	mer in						
35		f Composition:																				wane in Compositio	-Having Vinyl Homo					C Friction Coefficient (with same material) we Worm Amount (with same material) appect Strength After Weather Resistance	
40		part) o																				anosilo	1 Group		23°C -30°C			ficlent (vith s After W	
45		Blending Composition (part) of Composition:	Component (A)	100		Component (B)		Component (C)	1		Component (n)		Nylon 4, 6			DAG	100	Š	Polyarylate	121	PPE2	Amount (%) of Polyorganosiloxane in Composition	Amount (%) of carboxyl Group-Having Vinyl Honomer in Composition	Evaluation Result:	(kgf.cm/cm) (kgf.cm/cm) HPR (g/10 min.) (Temp.	DOP Brake Fluid	Gloss (%) Weld Strength Retentivity	Kinetic Priction Coeffici Relative Worn Amount (vil Izod Impact Strength Afte	-
50		Blenc	Co	77.	÷	Con		Š	b	55	Ü	×	ž	PBT	Z	2	2	Sad	8	2	ä	Amo	Com	Evalu	2 2 2 2	508	Mel	Re I	Tes

5	Example Example 47	9111 811	01	240)
15	Example Example 45	10 10 10 10 10 10 10 10 10 10 10 10 10 1	011 170111111	0.3 0.3 113 (240) 13 0.09 0.09 165
20	Example 14	01 1 0 0 1	011 110011111111	0.3 0.3 0.0 0.0 0.0 0.0 0.0 0.0
25	Example 43	1 1 101 1 10	114 101111111111	0.3 0.3 11 17 17 17 10 00 00 10 10 11
25	Example	1 01 1 0		0.3 111 145 (300) 00 00 16x10 <sup>-3</sup> 18
30	Example Example	· · ·	119 181111111111	0.3 0.3 0.9 0.09 1.6×10-3
35	·		:	1. Konomer in meterial) lance
40	Compositions		:	nosiscens in Composition Group-Having Vinyl Monomer 23°C 
45	(part) of		*	yorganosilox boxyl Group- gth; 23°C es30°C es30°C es30°C cs30°C es30°C
50	Composition (part) of Composition: A-1	A-2 A-4 A-4 A-4 B-1 B-2 B-3 Companent (C)	-1, -1, -1, -1, -1, -1, -1, -1, -1, -1,	Moment (i) of polyoginasitosme in Composition Composition Evaluation Result: Tead index Strength; 23°C (Medicalian) (Teap.) Composition Co
55	Blending Compon.	A-2 A-4 Component ( B-1 B-1 Component	C-1 C-2 C-3 C-3 Component Mylon 6 Mylon 6 Mylon 9 PVC PVC PVC PVC PVC PVC PVC PVC PVC PVC	Amount (% Campositi) Compositi) Evaluation Isod Imposition

10 15	CA) Example Example Example Example Example   A	10 10 10 10 10 10 10 10 10 10 10 10 10 1	25 25 25 25 25 25 25 25 25 25 25 25 25 2	7.6 16 16 16 20 0.3 0.3 0.3 1.5	22 (240) 29 (240) 13 (240) 25
25	Example Ex	01 01 08		0.3	28 21 (240) 1.0 1.0 88 92 92 16×10-3
35				Amount (1) of Polyorganosilosane in Composition Composition alustion Result:	The displayer Strength, 33°C (Med. of Community of Commun
40	(part) of (		· .	organosilom oxyl Group-1	rength; 23°C 1 (Temp.) tance: tance: Ratentivity (%) Racentivity (%) Amount (with samength After West
<b>45</b> 60	Blending Composition (part) of Composition: A-inner(A)	A-2 A-3 A-2 B-2 B-3 Component (C) C-1 C-2 C-3 Commonent (n)		Amount (%) of Polyorganosilozane in Composition Amount (%) of carboxyl Group-Having Vinyl Honom Composition Result:	lind impact Strangth) (Ref. officerin) (Temp. (Ref. officerin) (Temp. (Dop. and Resistance: Break Pluid (Sloss (R.) (Text. officerin) (Rinelin Friction Coeff (Relative Worn Mount)

35 40 	30	25		20	15	1	
<b>ਜ</b> '	TABLE 14 (Cont'd)	cont'd)				a	5
ition (part) of Composition:	Example 54	Example 55	Example 56	Example 57	Example 58	Example 59	Example 60
A-1 Component (B) B-1 B-2	g · · ·	0, 1,	or ' '	ō ' ' '	01'	g''	91 '
B-3 Component (c) C-3	g''	g''	30	g''	· g''	· gʻ'	- g: '
2-5 6-3	91	or '	10	10	g '	2'	10
Component (D) Nylon 6 Nylon 4, 6 Par							
PVC PVC HOPP PCC PCC PCC PCC PCC PCC PCC PCC PCC	1811	ı ' g '	8				
pps Pplyarylate PPE1 PPE2	uji.	1111		0111	1911	1 1 05 1	02
Amount (%) of Polyorganosiloxane in Composition	1.6	16	16	16	76	16	16
Amount (%) of Carboxyl Group-Having Vinyl Honomer in Composition		0.3	0.3	6.3	0.3	0.3	0.3
Evaluation Result:							
ised maper Extendibly 33°C (Mid-maper Extendibly 20°C) (Mid-man) (Yesp.) 10°C (Mid-man) (Yesp.) 10°C (Mid-man) (Yesp.) 10°C (Mid-man) (Yesp.) 10°C (Mid-man) (Yesp.) (	18 12 (200) 1.2 1.2 75 0.09 20x10-3	24 111 (220) 0 0 0 0 85 85 81 10.09 18×10 <sup>-3</sup>	18 10 0 0 0 0 176 82 82 13×10 <sup>-3</sup> 17	20 11 13 (300) 0 0 15 10 19	25 30 (300) 0 0 0 15 16 16×10·3 24	23 11 34 (260) 0 0 16×10-3 22	30 14 0 22 0 0 0 0.09 16x10 <sup>-3</sup> 29

While the invention has been described in detail and with reference to specific embodiments thereof, it will be apparent to one skilled in the art that various changes and modifications can be made therein without departing from the spirit and scope thereof.

#### Claime

1. A polyorganositoxane series thermoplastic resin compnsing a graft copolymer (V) obtained by graft polymerizing at least one kind of a vinyl monomer (IV) to a modified polyorganosiloxane (III) obtained by condensing from 90 to 99.8% by weight of an organosiloxane (I) having a structural unit represented by the following formula (A)

R'aSiO,4-a,2 (A)

wherein R represents a substituted or unsubstituted monovalent hydrocarbon group and n represents 0 or an integer of from 1 to 3, and from 10 to 0.2% by weight of a graft crosslinking agent (II) having both an unsaturated group represented by the following formula (B)

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$$CH_2 = C$$
(B)

wherein R2 represents a hydrogen atom or an alkyl group having from 1 to 6 carbon atoms, and an 25 alkoxysilyl group.

- 2. The polyorganosiloxane series thermoplastic resin as in claim 1, wherein 0.02 to 10% of R' in the formula (A) is a group containing a UV absorbing group.
- 3. The polyorganosiloxane series thermoplastic resin as in claim 1, wherein the residual group of R' in the formula (A) is at least one group selected from a methyl group, an ethyl group, a propyl group, a vinyl group, a phenyl group and those groups substituted with a halogen atom or a cyano group.
  - 4. The polyorganosiloxane series thermoplastic resin as in claim 1, wherein the residual group of R' in the formula (A) is at least one group selected from a methyl group, a vinyl group and a phenyl group.
  - 5. The polyorganosiloxane series thermoplastic resin as in claim 2, wherein the group containing the UV absorbing group has a structure represented by the following formula (II)

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wherein X represents

wherein R<sup>2</sup> represents an alkyl group having from 1 to 18 carbon atoms, Y represents H or OH, Z represents H OH and a represents H of the carbon atoms, Y represents H of the represents H, OH or OR2 wherein R2 is the same as defined above, and n represents an integer of from 1 to 5. with coulded the control of the same as defined above, and n represents an integer of from 1 to 5. 55 5, with proviso that when Y is H, at least one Z is OH.

6. The polyorganosiloxane series thermoplastic resin as in claim 1, wherein the graft crosslinking agent (II) is at least one member selected from p-vinylphenylmethyldimethoxysilane, 1-(m-vinylphenyl)-methyldimethoxysilane, 3-(p-vinylphenyxy)-methyldimethoxysilane, 3-(p-vinylphenyxy)-methyldimethoxysil methyldimethylisopropxysilane, 2-(p-vinylphenyl)ethylmethyldimethoxysilane,

propylmethyldiaethoxysilane. 3-(p-viny/benzoyloxy)propylmethyldimethoxysilane. 1-(o-vinylphenyl)-1.1.2-timethyl-2.2-dimethoxydsilane. 1-(p-vinylphenyl)-1.1-diphenyl-3.3-diethoxydsiloxane, m-vinylphenyl-1-(methoxysily)propyldiphenylsilane and [3-(P-isopropenylbenzoylamino)propyll-phenyldiphenyl-2-

- 7. The polyograpisaloxane series thermoplastic resin as in claim 1, wherein the modified polyograpisaloxane (III) is obtained by polyoridensing the organosiloxane (II) having the structural unit represented by the following formula (A)
- R2nSiO(4n)2 (A)
- wheren RP represents a substituted or unsubstituted monovalent hydrocarbon group and n represents O or to an integer of from I to 3, an organic silicone compound (III) having both the group containing the reactive unsaturated group and an alkoxy group, and an organic silicone compound (IVI) having both a group containing a IVV absorbing group and an alkoxy group in the presence of an emulsilying agent.
- The polyorganosibxane senes thermoplastic resin as in claim 7, wherein the emulsilying agent is at least one member selected from an aliphatic substituted benzenesullonic acid, an aliphatic substituted of aphthalenesullonic acid, an aliphatic substituted diphenyl ether sulfonic acid, each aliphatic substituted diphenyl ether sulfonic acid, each aliphatic group having a carbon chain length of from 6 to 18 carbon atoms.
  - The polyorganosiloxane series thermoplastic resin as in claim 1, wherein the silicon atom number in the modified polyorganosiloxane (III) is from 100 to 10,000.
  - The polyorganosiloxane senes thermoplastic resin as in claim 1, wherein the polystyrene calculated weight average molecular weight of the modified polyorganosiloxane (III) is from 10,000 to 5,000,000.
- 11. The polyorganosiloxane series thermoplastic resin as in claim 1, wherein the vinyl monomer (IV) is at least one member selected from styrene, a-methylstyrene, sodium styrenesullonate, methyl meth acrylate, ethyl methacrylate, atly methacrylate, atly methacrylate, are methacrylate, methyl acrylate, ethyl acrylate, butyl acrylate, butyl methacrylate, dimethylaminoesthyl methacrylate, acrylotic, ethyl acrylate, butyl acrylate, soprene, chloroprene, vinyl acetate, vinyl chloride, vinylidere chloride, trialtyl isocyanurate, acrylic acid, methacrylate, acid, hypenylmaleimide, N-cyclohexylmaleimide, maleic anhydride, crotonic acid, cinnamic acid, itaconic acid, maleic acid, glycidyl méthacrylate, glycidyl acrylate, vinylglycidyl ether, allylglycidyl ether, glycidyl ether of hydroxyalkyl (meth)-acrylate, glycidyl ether of polyalkylene glycol (meth)-acrylate, and glycidyl itaconate.
  - 12. The polyorganosiloxane series thermoplastic restn as in claim 1, wherein the vinyl monomer (IV) comprises at least one member selected from a carboxyl group-containing vinyl monomer and an epoxy group-containing vinyl monomer, and other vinyl monomer.
- 13. The polyorganositoxane series thermoplastic resin as In claim 1, wherein the graft ratio of the graft copolymer (V) is at least 20% by weight.
  - 14. The polyorganosiloxane series thermoplastic resin as in claim 1, wherein the graft ratio of the graft copolymer (V) is at least 80% by weight.
  - 15. A thermoplastic resin composition comprising from 1 to 99% by weight of the polyorganosiloxane
- series thermoplastic resin of claim 1 and from 99 to 1% by weight of other polymer. 16. The thermoplastic resin composition as in claim 15, wherein the other polymer is at least one polymer selected from polybutadiene, a butadiene-styrene copolymer, an acrylonitrile-butadiene copolymer, polyisoprene, natural rubber, acryl rubber, an ethylene-propylene copolymer, an ethylene-propylene-diene copolymer, chlorinated butyl rubber, chlorinated polyethylene, a styrene-butadiene block copolymer, a styrene-butadiene-styrene block copolymer, a styrene-butadiene-styrene radialtereblock copolymer, a 45 hydride of the above block copolymer, polypropylene, polyethylene, polystyrene, a styrene-acrylonitrile copolymer, rubber-reinforced, polystyrene (HIPS), an acrylonitrile-butadidne-styrene resin (ABS resin), an acrylonitrie-ethylenepropylene-styrene resin (AES resin), a methyl methacrylate-butadiene-styrene resin (MBS resin), an acrylonitrile-butadiene-methyl methacrylate-styrene resin, an acrylonitrile-n-butyl acrylatestyrene resin (AAS resin), polyvinyl chloride, polycarbonate, polyethylene terephthalate, polybutylene terephthalate, polyacetal, polyamide, an epoxy resin, polyvinylidene fluoride, polysulfone, an ethylene-vinyl acetate copolymer, a PPS resin. polyether ether ketone, a PPO resin, polyacrylate, a styrene-methyl methacrylate copolymer, a styrene-maleic anhydride copolymer, a rubber-modified PPO resin, a styrenemaleimide series copolymer a rubber-modified styrene-maleimide series copolymer a polyamide series elastomer, and a polyester series elastomer.
  - 17. A thermoplastic resin composition comprising
  - (A) from 5 to 90% by weight of a graft copolymer obtained by graft polymerizing from 95 to 10% by weight of the sum of a carboxyl group—containing vinyl monomer and other vinylic monomer to from 5 to 90% by weight of the modified polyborganosiloxane (III) of claim 1.

- (8) from 0 to 80% by weight of a graft copolymer obtained by graft polymerizing from 95 to 10% by weight of other verytic monomer than the carboxyl group-containing viryl monomer to from 5 to 90% by weight of the modified polyograpoisosane copolymer.
  - (C) from 0 to 80% by weight of a styrene series resin, and
- (D) from 10 to 90% by weight of at least one thermoplastic resin selected from a polyamide senes resin, a polyester senes resin, a polyer/conate senes resin, a winyl chloride senes resin, an olefinic resin, a polyacetal senes resin, polyphenylene sulfide series resin, a polyarylate senes resin, and a polyphenylene ether senes resin
- 18. The thermoplastic resin composition as in claim 17, wherein the carboxyl group-containing viryl monomer is at least one member selected from acrylic acid, methacrylic acid, crotonic acid, cinnamic acid, itaconic acid, maleic acid, and maleic anitydride.
  - 19. A thermoplastic resin composition comprising

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- (A) from 90 to 10% by weight of a graft copolymer obtained by graft polymerizing from 95 to 10% by weight of the sum of an epoxy group-containing vinyll monomer and other vinyl monomer to from 5 to 90% by weight of the modified polyroganosioxane (III) of claim 1, and
  - (B) from 10 to 90% by weight of at least one thermoplastic resin selected from a polyphenylenesulfide senes resin, a polyamide senes resin, a polyester series resin, a polycarbonate series resin, a vinyl chloride senes resin, an olefinic resin, a polyacetal series resin, a polyarylate series resin, and a polyphenylene ether senes resin.
  - 20. The thermoplastic resin composition as in claim 19, wherein the epoxy group-containing vinyl monomer is at least one monomer selected from glycidyl methacrylate, glycidyl acrylate, vinylglycidyl ether, glycidydyl ether, glycidyd ether, glychyd ether,

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